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2	Characterization of chlorate reduction in the haloarchaeon Haloferax						
3	mediterranei.						
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15							
16	Abbreviations: DT, dithionite; MV, methylviologen; NarGH, respiratory nitrate						
17	reductase; PCRB, perchlorate-respiring bacteria; CRB, chlorate-respiring						
18	bacteria.						
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2 Abstract

Background: *Haloferax mediterranei* is a denitrifying haloarchaeon using nitrate
as a respiratory electron acceptor under anaerobic conditions in a reaction
catalysed by pNarGH. Other ions such as bromate, perchlorate and chlorate can
also be reduced.

7 Methods: *Hfx. mediterranei* cells were grown anaerobically with nitrate as 8 electron acceptor and chlorate reductase activity measured in whole cells and 9 purified nitrate reductase.

10 Results: No genes encoding (per)chlorate reductases have been detected either 11 in the Hfx. mediterranei genome or in other haloarchaea. However, a gene 12 encoding a chlorite dismutase that is predicted to be exported across the 13 cytoplasmic membrane has been identified in *Hfx. mediterranei* genome. Cells 14 did not grow anaerobically in presence of chlorate as the unique electron 15 acceptor. However, cells anaerobically grown with nitrate and then transferred to chlorate-containing growth medium can grow a few generations. Chlorate 16 17 reduction by the whole cells, as well as by pure pNarGH, has been characterised.

18 No clear chlorite dismutase activity could be detected.

19 Conclusions: *Hfx. mediterranei* pNarGH has its active site on the outer-face of 20 the cytoplasmic membrane and reacts with chlorate and perchlorate. Biochemical 21 characterisation of this enzymatic activity suggests that *Hfx. mediterranei* or its 22 pure pNarGH could be of great interest for wastewater treatments or to better 23 understand biological chlorate reduction in early Earth or Martian environments.

General significance: some archaea species reduce (per)chlorate. However, results here presented as well as those recently reported by Liebensteiner and co-workers [1] suggest that complete perchlorate reduction in archaea follows different rules in terms of biological reactions. Keywords: Halophile; Archaeon; respiratory nitrate reductase; chlorate reduction; anaerobic respiration; chlorite dismutase.

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3 **1. Introduction**

4 During the last 10 years, perchlorate (ClO_4) and chlorate (ClO_3) have been detected in several water supplies, ground waters, agricultural crops and 5 6 even in soils as a result of human activities [2]. Perchlorate is used in the 7 manufacture of propellants, explosives and pyrotechnic devices [3]. Perchlorate 8 salts are extremely soluble, non-volatile, non-reactive and chemically very stable. 9 The high water solubility and poor adsorption of perchlorate to soil and organic 10 matter make its high mobility in the environment possible [4]. The concerns about 11 perchlorate toxicity are its interference with iodide uptake by the thyroid gland, 12 and the related potential carcinogenic effects [5]. Chlorate is present in several 13 herbicides and defoliants, and it is released when chlorine dioxide (CIO_2) is used as a bleaching agent in the paper and pulp industry [6]. In humans, CIO_3^{-1} may 14 15 cause thyroid lesions and anaemia. Because of these health concerns, several organizations such as the World Health Organization or the Environmental 16 17 Protection Agency have advised that (per)chlorate in water intended for human 18 consumption should be minimized [7].

Perchlorate and chlorate are ideal electron acceptors for microorganisms due to their high redox potentials ($CIO_4^{-7}/CI^{-}E_o=1.287$ V; $CIO_3^{-7}/CI^{-}E_o=1.03$ V) [8]. It has been proposed that in perchlorate-respiring bacteria (PCRB) the (per)chlorate-reduction pathway consists of the (per)chlorate reductase, which sequentially reduces perchlorate to chlorate and in turn chlorate to chlorite (CIO_2^{-7}), via sequential two-electron transfers [6,7]. Finally, chlorite dismutase

transforms chlorite into chloride and oxygen [9-13]. Perchlorate-respiring bacteria (PCRB) are ubiquitous in the environment, and are mainly facultative anaerobes and denitrifiers [14,15]. Perchlorate reductases isolated from PCRB react with both perchlorate and chlorate [6], while chlorate reductases expressed by chlorate-respiring bacteria (CRB) do not reduce perchlorate [16]. It has also been demonstrated that perchlorate and chlorate reductases isolated from some PCRB recognize nitrate as substrate [17].

8 Nitrate is also often present in environments where perchlorate or chlorate 9 are faced as contaminants [18]. In the denitrification pathway, nitrate is sequentially reduced to dinitrogen gas: $NO_3^- \rightarrow NO_2^- \rightarrow NO \rightarrow N_2O \rightarrow N_2$ [19]. 10 Several nitrate reductases involved in anaerobic nitrate reduction also reduce 11 chlorate. However, these nitrate reductases have active sites facing the 12 13 membrane potential negative side (nNars) and the nitrate transporters that 14 deliver nitrate into the cytoplasm do not recognize chlorate, thus preventing the 15 potentially damaging intracellular reduction of chlorate to cytotoxic chlorite. In 16 previous studies on the respiratory nitrate reductase (NarGH) from Haloferax 17 mediterranei, a denitrifying halophilic archaeon able to use nitrate as nitrogen source for growth or as electron acceptor under anaerobic conditions [20-22], it 18 was demonstrated that this enzyme has an active site facing the membrane 19 20 potential positive face (pNars) and is able to reduce chlorate [18]. The extra-21 cytoplasmic active site could be accessible to chlorate and so, this reaction might 22 take place in the environment leading to the question of whether it could support 23 growth and also whether it could reduce perchlorate, a substrate for which the

role as electron acceptor in the reactions catalyzed by nitrate reductases has
 been poorly described in haloarchaea.

3 Recent results reveal that (per)chlorate reductases establish a distinct group with the archaeal p-type NarG nitrate reductases as the closest relatives 4 into dimethyl sulfoxide (DMSO) reductase family [23]. It has also been proposed 5 6 that chlorate reduction was built multiple times from type II dimethyl sulfoxide 7 (DMSO) reductases and chlorite dismutases [24, 25]. This work summarises the 8 biochemical characterisation of the NarGH chlorate reductase activity and 9 discusses different strategies that might be used by haloarchaea to deal with the 10 chlorite produced during chlorate reduction.

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12 **2. Materials and Methods**

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14 **2.1 Strains, media and growth conditions**

15 Hfx. mediterranei (ATCC 33500T) was grown anaerobically with nitrate (100 mM) as electron acceptor as previously described [18], in a 25% (wt/vol) 16 17 mixture of salts (25% SW) [26] and 0.5% yeast extract (complex media). Cultures 18 with chlorate (100mM) as electron acceptor were prepared in the same way. 19 Growth was monitored by measuring the optical density at 600 nm. In some 20 experiments, *Hfx. mediterranei* cells were grown as already described to induce 21 the denitrification pathway (nitrate as electron acceptor). After that, cells were 22 harvested by centrifugation at 15000 x g for 20 min at 4 °C in a Beckman J2-21 23 centrifuge, washed with 25% SW, centrifuged again at 15000 x g for 20 min at 4 °C and transferred to fresh anaerobic complex medium (25% SW and 0.5% yeast
 extract) containing chlorate at different concentrations ranging from 5 to 100 mM.
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5 **2.2** Purification of respiratory nitrate reductase and characterization of the

6 NarGH chlorate reductase activity

7 All the purification steps were carried out at 25 °C following the protocol 8 previously described [20]. The chlorate reductase activity of the NarGH was 9 measured using two different methods: i) colorimetric o-toluidine assay, which 10 allows chlorate quantification [27] and ii) methylviologen method (substrate-11 dependent oxidation of reduced methylviologen) [28]. For the o-toluidine assay, 12 the reaction mixture contained, in a final volume of 1ml: 50 mM Tris-HCl pH (7-9), 13 0-2M NaCl or KCl, 5 mM MV (electron donor), 50 mM KClO₃ (substrate), 10 mM 14 $Na_2S_2O_4$ (freshly prepared in 0.1 M NaHCO₃) and 40 µl of pure enzyme (final 15 protein concentration around 0.03 mg protein per mL was constantly present in 16 the reaction mixture). The enzymatic activity was tested at temperatures between 17 25 and 70 °C, but most of the assays were developed at 35 °C. After 5 min of 18 incubation to allow enzymatic reaction, 0.25 ml of o-toluidine (0.4 g/l) and 1.25 ml of concentrated HCI were added to reaction mixture. O-toluidine and HCI 19 20 additions destroy the protein and as a consequence, the enzymatic reaction is 21 stopped. The absorbance related to the yellow hologuinone finally produced in 22 the colorimetric reaction was checked at 490 nm. NarGH chlorate reductase 23 activity is expressed as micromoles of KCIO₃ reduced per minute and chlorate 24 reductase specific activity is expressed as micromoles KCIO₃ reduced per minute

1 per milligram of protein. All the assays were carried out in triplicate and against a 2 control assay without enzyme. The control without enzyme was used for two 3 different purposes: i) to check that there is no chlorate reduction when removing 4 the enzyme and ii) to quantify chlorate concentration within the reaction mixture 5 at zero time. This ensures that the kinetics of chlorate reduction take into account 6 the chlorate concentration within the reaction mixture at zero time. To determine 7 the optimal pH for chlorate reductase activity, 50 mM MES (pH 5.5.-6.7), 50 mM 8 MOPS (pH 6.5-7.9) and 50 mM carbonate (pH 9-11) buffers were also prepared 9 containing the aforementioned reaction mixtures. For the methylviologen assay, 10 chlorate reductase activity was measured spectrophotometrically in quartz 11 cuvettes equipped with rubber septa by monitoring the oxidation of reduced MV $(\epsilon_{600 \text{ nm}} = 13700 \text{ M}^{-1} \text{ cm}^{-1})$ in presence of chlorate at different temperatures [28]. 12 13 The reaction mixture contained, in a final volume of 800 µl, 50 mM Tris buffer pH 14 8, 0-2M NaCl or KCl, 5 mM MV and an appropriate amount of pure enzyme (final protein concentration around 0.03 mg protein per mL was constantly present in 15 16 the reaction mixture). The assay mixture was flushed with nitrogen for 10 min, 17 and 5 mM dithionite solution (degassed and freshly prepared in 0.1 M NaHCO₃) 18 was added until an absorbance of 3.0 at 600 nm was obtained. The reactions, 19 incubated for 1 min at 40 °C, were initiated by the addition of KCIO₃ (nitrogen 20 flushed) to a final concentration between 0-50 mM. Alternative electron acceptors 21 were tested in the same assay system, except that chlorate was replaced by CIO_4^- , NO_3^- , IO_3^- , BrO_3^- and SeO_4^{2-} (potassium salts). All the assays were 22 23 carried out in triplicate and against controls without enzyme or without the 24 electron acceptors.

1 The MV assay method was also used to follow the chlorate reduction 2 using whole cells previously grown with nitrate or chlorate as electron acceptors. 3 In this case, harvested cells were resuspended in 50 mM Tris buffer containing 0.5 M NaCl up to a final O.D. around 0.2. The reaction mixture (1.2 ml final 4 volume) contained 1 ml resuspended cells, 5 mM MV, 5 mM dithionite solution 5 6 and 0-25 mM substrate. The assay was developed at room temperature. All the 7 assays were carried out in triplicate and against controls without cells or without 8 the electron acceptors. Data obtained by MV method were processed using the 9 Michaelis–Menten equation. The values of V_{max} and K_m were determined by 10 nonlinear regression analysis of the corresponding Michaelis-Menten curves 11 (rate vs. [CIO₃]) using the algorithm of Marguartd–Levenberg with the SigmaPlot 12 program (Jandel Scientific, version 1.02). The protein content was determined by 13 the Bradford method, with bovine serum albumin (fraction V) as a standard.

14 In order to check the effect of other anions (e.g. bromate and (per)chlorate) on nitrate reduction catalyzed by *Hfx. mediterranei* pNarGH, the 15 nitrate reduction was also measured as previously described [20]. In that 16 17 instances, the standard reaction mixture contained 4mM MV (artificial electron 18 donor), 18 mM KNO₃, and different chlorate, perchlorate and bromate 19 concentrations (from 0 up to 18 mM). We followed the nitrite production by pNar 20 using Griess method [29]. Nitrate reductase specific activity is expressed as 21 micromoles of NO₂ appearing per minute per milligram of protein [20].

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4 **3. Results**

5 **3.1 Chlorate reduction by** *Hfx. mediterranei* cells

6 In order to analyze the capacity of Hfx. mediterranei cells to reduce 7 chlorate, the cell growth was checked in anaerobic media using nitrate or 8 chlorate as electron acceptors. When nitrate is present within the anaerobic 9 media, denitrification is induced and as a consequence, *Hfx. mediterranei* is able 10 to use nitrate as electron acceptor to support growth, as previously described 11 [19-22, 30]. However, no growth was observed if chlorate was added as the 12 unique electron acceptor to the anaerobic media. Specific growth rate (μ) and cell doubling time (t_d) for cultures under each of the assayed conditions are 13 14 summarised in table 1. These results suggest that this haloarchaeon is unable to 15 express chlorate inducible genes coding for a (per)chlorate reductase system. Recently, the *Hfx. mediterranei* genome draft has been obtained and no genes 16 17 encoding (per)chlorate reductases have been detected 18 (www.ncbi.nlm.nih.gov/genome/?term=haloferax+mediterranei).

However, when nitrate-respiring cells (anoxic conditions) were harvested at the beginning of the stationary phase of growth, washed and transferred to a fresh anaerobic chlorate media, the cells were able to use chlorate as electron acceptor. In this case, the optical density reached at stationary phase of growth is 0.8 (after 150 hours of batch culture), t_d is 27 hours and a μ of 0.019 h⁻¹ is achieved. When the cells pre-grown with nitrate were transferred into chlorate1 containing medium more than twice, cells were not able to grow anymore. This 2 suggests that pre-induced respiratory nitrate reductase could be involved 3 physiologically in chlorate reduction supporting limited cell culture growth. In earlier studies, it was stated that (per)chlorate and nitrate reduction were 4 catalyzed by the same enzyme (a nitrate reductase) in bacteria [31]. This implies 5 6 that the inability of several denitrifiers to grow using (per)chlorate is due to: (1) 7 failure to induced nitrate reductase in the presence of chlorate alone and (2) the 8 toxicity chlorite produced by nitrate reductase when chlorite dismutase is absent.

9 This explanation is supported by experiments where chlorate, perchlorate and 10 bromate reduction were measured using whole *Hfx. mediterranei* cells previously 11 grown with nitrate as electron acceptor (Fig. 1A), where the order of the reaction 12 velocities was $CIO_3^- > NO_3^- > BrO_3^- > CIO_4^-$, while no reduction activity was 13 detected in presence of IO_3^- or SeO_4^- , similarly to the purified enzyme (Fig. 1B). 14 This characteristic could be the explanation for the *Hfx. mediterranei* NarGH 15 chlorate reductase activity.

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17 **3.2** Characterization of the NarGH chlorate reductase activity

A preliminary report that pNarGH could reduce chlorate has previously been made [32]. A more detailed study was undertaken of this reaction in the present study to explore the use of this enzyme (and/or the whole cells) in waste water treatments. The capacity of pure NarGH to reduce substrates such as iodate, selenate, bromate and perchlorate has been checked. The results obtained (Fig. 1B) have been compared with the nitrate reduction. Consistent with the results from the whole cell experiments, no activity was detected when

1 iodate or selenate were used as substrate. The absence of NarGH reactivity 2 towards selenate was previously pointed out and it has been proposed that this 3 fact could be taken into account to distinguish pNars from nNars (the latter type 4 of Nars are highly reactive towards selenate) [22]. Nevertheless, pure Hfx. 5 mediterranei NarGH reduced chlorate, bromate and perchlorate with reaction velocities in the order of 0.25 µmol min⁻¹ mg⁻¹, 0.15 µmol min⁻¹ mg⁻¹ and 0.11 6 μ mol min⁻¹ mg⁻¹, respectively. The order of effectiveness was ClO₃ > NO₃ > 7 $BrO_3 > ClO_4$ which is the same pattern observed from the activity 8 9 measurements using whole cells (Fig. 1A). This pattern correlates with that 10 observed in whole cells suggesting it is due to the NarGH present in the 11 membranes of these cells.

12 The pH dependence of enzyme activity in the range 5 to 11 revealed that the optimal pH for pNarGH chlorate reductase activity was around pH 8 at 40 °C 13 14 (which is the average temperature detected in the Hfx. mediterranei natural environment) (Fig. 2A). pNarGH nitrate reductase activity also showed maximum 15 activity at pH values around 8 [20]. Bacterial chlorate reductases recently 16 17 described had a temperature optimum between 40 and 70°C, which is perhaps 18 unexpected because most bacteria from which they were purified are mesophilic 19 [16].

20 NarGH chlorate reductase activity was also measured at different NaCl or 21 KCl concentrations at 40 °C (Fig. 2B). In those reaction mixtures containing NaCl 22 the enzymatic activity increased when the NaCl concentrations were increased 23 up to 1.4 M and at higher NaCl concentrations the activity remained stable. 24 However, in those assays carried out in presence of KCl, the enzymatic activity

increased when the KCI concentrations were increased up to 1M, but at higher KCI concentrations, a decrease in the specific enzymatic activity could be detected. This result could be related to the fact that in pNarGH the catalytic subunit is oriented to the positive side of the membrane [22] and in natural saltmarsh environments, NaCI is the predominant salt (instead of KCI which is predominant in the cytoplasm).

7 In another set of experiments, NarGH chlorate reductase activity was measured using an assay mixture with different NaCl concentrations at 8 9 temperatures from 25° to 70 °C (Fig. 3). The maximum specific chlorate 10 reductase activity was detected in the presence of low NaCl concentrations at 35 11 ^oC. At salt concentrations higher than 0.4 M, the maximum specific activity could 12 be detected at higher temperatures (between 40-45 °C). The data was subject to an Arrhenius analysis that revealed that the lowest activation energy (around 2.2 13 ± 0.2 J mol⁻¹ in presence of 0.8-1.6 M NaCl) was observed at the highest NaCl 14 15 concentrations (versus 5 ± 0.5 J mol⁻¹ in presence of 0-0.4 M NaCl. These results strongly support the fact that activation energy of a halophilic enzyme is lower at 16 17 high salt concentrations. However, the higher the salt concentrations, the lower 18 NarGH chlorate reductase activity was observed. This is not the pattern expected from halophilic enzymes, which are characterised by high enzymatic activity 19 20 values at high salt concentrations. The pattern here described for NarGH chlorate 21 reductase activity differs from the behaviour previously described for NarGH nitrate reductase activity. In the latter, nitrate reduction was not strongly 22 23 dependent on temperature at different NaCl concentrations [20].

1 Like other halophilic nitrate reductases from the genus Haloferax, NarGH 2 from *Hfx. mediterranei* presented a remarkable thermophilicity and worked well up to 70 °C with nitrate as substrate and this activity did not show a direct 3 dependence on salt concentration [20]. However, working with chlorate as 4 5 substrate, NarGH exhibited higher specific activity at low temperatures in 6 presence of low salt concentrations (Fig. 3). This may reflect the different 7 molecular geometry of the substrates (planar triangular for nitrate, triangular 8 pyramidal for chlorate or tetragonal pyramidal for perchlorate).

9 Kinetic parameters of NarGH were determined using different concentrations of chlorate (as substrate), in the presence of 50 mM Tris buffer 10 11 (pH 8.0) containing 0.2 M NaCI. The halophilic enzyme followed Michaelis-12 Menten kinetics. Apparent V_{max} and K_m values for chlorate were 0.280 \pm 0.003 μ mol min⁻¹ mg prot.⁻¹ and 2.41 ± 0.16 mM, respectively. These values are higher 13 than those described in several chlorate reductases isolated from bacteria such 14 15 as *Pseudomonas chloritidismutans* [16], which is in agreement with the fact that although NarGH is able to reduce chlorate, it is not its natural substrate. The 16 17 turnover numbers as well as the specificity constant have been obtained taking 18 into account the molecular mass of the NarGH isolated from Hfx. mediterranei [20] and the V_{max} obtained from reactions where chlorate or nitrate act as 19 20 substrates; results are summarized in table 2.

Finally, nitrate reduction activity has been measured in presence of different anions such as perchlorate, chlorate and bromate using pNarGH from *Hfx. mediterranei* or the whole cells. As previously mentioned, it is quite common to find environmental samples or waste water samples containing nitrate and

1 (per)chlorate. We expected to see competitive inhibition of nitrate reduction due 2 to the presence of (per)chlorate ions in the reaction mixture. That's why, we 3 hypothesised that when the highest oxyanion concentration was added, the lowest nitrite concentration would be produced. However, the opposite effect was 4 5 observed: the nitrite production was even more extensive when bromate, chlorate 6 and perchlorate were present within the reaction mixture containing nitrate as 7 substrate (Fig. 4). From these results, we conclude that in the presence of nitrate, 8 these oxyanions are neither inhibitors nor alternative substrates. We suspect that 9 the high redox potential defining each of those chemical compounds has an 10 important role in this enzymatic mechanism. Thus, they could act as electron 11 carriers in the reduction of nitrate to produce nitrite. Another interesting feature to 12 highlight from these results is that the reaction in presence of nitrate + other 13 oxyanion requires more time to take effect (nitrite is not detected prior to 2-3 14 minutes of incubation), but when nitrite production starts, the reaction is quicker 15 than that taking place in absence of bromate, chlorate and perchlorate. These results are relevant when trying to explore the feasibility of bioremediation of 16 17 waste water samples containing more than one of those compounds.

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19 **4. Discussion**

The results reported here indicate that *Hfx. mediterranei* is able to use (per)chlorate as final electron acceptor in absence of oxygen using the nitrate reductase (pNarGH) involved in denitrification process. Some recent reports support the existence of separate pathways for (per)chlorate and nitrate reduction under anaerobic conditions, although they have not completely

eliminated the potential of shared enzymes, being used for (per)chlorate and
nitrate reduction in some bacteria [33]. The physiological study previously
discussed suggests that chlorate respiratory enzymes are not inducible in *Hfx. mediterranei* by chlorate under anaerobic conditions, which make sense taking
into account that genes coding for (per)chlorate reductases have not been
identify in *Hfx. mediterranei* genome.

7 One interesting aspect to be pointed out is what happens to the chlorite 8 produced by NarGH during chlorate reduction by *Hfx. mediterranei*. The analysis 9 of the *Hfx. mediterranei* genome draft shows that the gene encoding a putative 10 chlorite dismutase is present (Fig. 5), while no genes coding for (per)chlorate 11 reductases have been detected either in the Hfx. mediterranei genome nor in 12 haloarchaea Database searches other up to now. (HALOLEX: 13 www.halolex.mpg.de/public/) have pointed out that chlorite dismutase gene from Hfx. mediterranei is a homolog to pitA from Hfx. volcanii, which is a fusion 14 15 between chlorite dismutase-like and antibiotic biosynthesis monooxygenase-like domains within a single open reading frame. This fusion has been also described 16 17 from other haloarchaea and may represent a modification to limited oxygen 18 availability [34]. Preliminary studies to detect chlorite dismutase activity in Hfx. 19 mediterranei whole cells and extracts have been carried out in our laboratory 20 following oxygen production in presence of chlorite using Durham tubes and 21 oxygen electrodes. An activity toward chlorite could not clearly be identified either in whole cells or in cell extracts. Some recent work from some bacteria suggest 22 23 that some chlorate reduction genes might constitute transposons flanked by 24 insertion sequences which show the potential to move horizontally [24]. The

phylogenetic analysis carried out using bacterial genomes reveals that chlorate reduction was evolved multiple times from type II DMSO reductases and chlorite dismutases [24]. It has also been suggested that chlorite dismutase has been mobilised at least once from (per)chlorate reducers to build chlorate respiration [24]. More studies focused on that subject should be done in archaea to understand how chlorate respiration was built in this domain.

7 Other studies from hyperthermophilic archaea highlight that although 8 (per)chlorate can be used as electron acceptor using enzymes belonging to the 9 type II subgroup of DMSO reductase family, no chlorite dismutase activity has 10 been detected. In that case, the authors demonstrate that chlorite is eliminated 11 by interplay of abiotic and biotic redox reactions involving sulphur compounds 12 instead of being enzymatically split into chloride and oxygen [1]. This work has 13 been carried out using Archaeglobus fulgidus, which is a hyperthermophilic 14 archaeon that thrive in environments resembling those of early Earth.

In the study presented here, we have not added specific sulphur 15 compounds to check the mentioned interplay of abiotic and biotic redox reactions 16 17 and no evidence on that subject has been reported so far from haloarchaea. So, 18 it would be worth exploring these abiotic and biotic redox reactions involving 19 sulphur in the future within in member of the Halobacteriaceae family. Regarding 20 this, it is interesting to draw attention to: i) the natural salted water where these 21 microorganisms live contains sulphur salts, ii) when growing the haloarchaea in contains sulphur salts (ammonium sulphate, the lab, salted water also 22 23 magnesium sulphate, etc.) and iii) some of the protocols used to purify proteins 24 from haloarchaeal involve buffers containing sulphur compounds (i.e ammonium

sulphate). Therefore, we cannot dismiss the possibility that abiotic and biotic
redox reactions involving sulphur compounds take place in haloarchaea. Some
works related to bacterial communities in marine sediments have also stated
recently that there are (per)chlorate effects on metabolic pathways related to
sulphur [35].

This research as well as other recent published reports suggest that haloarchaea can reduce perchlorate and chlorate anaerobically. This fact has recently been connected to studies of the anaerobic Martian environment where perchlorate among the salts as detected by the Phoenix Lander on Mars, may support halophilic life in a similar way to the halophilic environments on Earth [36].

12 The results presented show that (per)chlorate anions can be efficiently 13 removed from the media where the *Hfx. mediterranei* cells are growing. 14 Perchlorate and chlorate reduction take place efficiently thanks to *Hfx.* 15 *mediterranei* cells, even in presence of low salt concentrations. This fact could 16 suggest that changes in the structure of the enzyme, and therefore the chlorate 17 reductase activity, could be caused by salt.

If this is the case for other denitrifying halophilic archaea, then those type of microorganisms could be excellent models for the bioremediation of brines [37]. Most of the ground waters or waste waters contaminated with chlorate also contain nitrate [16] and several previous studies have shown that (per)chlorate and nitrate are simultaneously degraded by several bacteria [38]. A better understanding of the factors that regulate the expression of the enzymes involved in (per)chlorate, chlorate, and nitrate reduction are important as several

1 microorganisms could play a key role in water bioremediation. The processes 2 based on bioremediation could replace or even improve those protocols where 3 perchlorate and nitrate are removed from brines by ion exchange techniques [5]. 4 Several studies have highlighted that (per)chlorate-reducing bacteria removed 5 (per)chlorate at such slow rates to make them impractical for application in 6 treatment systems. Moreover, the enzymes involved in this pathway are only 7 active at low salt concentrations [5]. Other previous studies reported rapid 8 perchlorate and nitrate removal using mixed cultures in presence of 10% NaCl 9 [39]. Other systems use permeable barriers containing vegetable oil that would 10 promote the degradation of (per)chlorate [7]. However, each process has a 11 limitation, namely salt tolerance.

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13 **5.** Conclusions

14 Respiratory nitrate reductases, mainly those belonging to the pNarGH 15 group [20] may have a relevant role in (per)chlorate reduction in those halophilic 16 archaea lacking genes coding for (per)chlorate reductases. Biological 17 per(chlorate) reduction by ancient archaea might have taken place during pre-18 anthropogenic times thanks to (per)chlorate reductases or even nitrate 19 reductases. As a consequence of these enzymatic activities members of the 20 archaea domain may have prevented perchlorate accumulation in early Earth 21 giving rise to the environmental conditions characterising the Earth [1]. Taking into account the biochemical parameters defining NarGH chlorate reductase 22 23 (reported here) and NarGH nitrate reductase activities in Hfx. mediterranei 24 [20,22], waste water treatment approaches could consider the relevance of

1 halophilic denitrifiers to explore a role in bioremediation in the near future. 2 Preliminary results obtained in our lab reveal that chlorate removal by Hfx. 3 mediterranei cells is more efficient than per(chlorate) removal. In those media containing 5 mM chlorate, the final chlorate concentration quantified after 150 4 hours of incubation was 0.2 mM. On the other hand, nitrate reduction is not 5 6 inhibited in presence of either (per)chlorate or bromate, it can even be concluded 7 that bromate is able to slightly stimulate nitrate rduction (see results summarised 8 in figure 4). So, the same microorganism could reduce nitrate and chlorate in 9 presence of other ions thanks to the nitrate reductase under microaerobic or 10 anaerobic conditions. These results are guite interesting in terms of waste water 11 bioremediation purposes because most of the waste water samples containing 12 nitrate also contain chlorate and other oxyanions. The removal ratio for chlorate 13 estimated in our study is around 4.8 mM chlorate in approximately 6 days. 14 Although the removal process is not really fast, the removed concentration is one 15 of the highest described at the time of writing when using microorganism for waste water bioremediation [40,41]. Furthermore, one of the advantages of using 16 17 *Hfx. mediterranei* cells or its pNarGH is that nitrate reduction is not inhibited in 18 presence of (per)chlorate (and vice versa) at high salt concentrations. These 19 results open a new way to explore bioremediation processes making 20 haloarchaeaon-based remediation of brines and waste waters feasible.

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23 Acknowledgements

1	This work was funded by research grant from the MINECO Spain
2	(CTM2013-43147-R), MEC Spain (BIO2008-00082), Generalitat Valenciana
3	(GV/2011/038) and University of Alicante (GRE09-25).
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Legend to figures Figure1. Reduction of different substrates by whole *Hfx. mediterranei* cells (A). Cells were grown under anaerobic conditions with nitrate as electron acceptor as cited in materials and methods. Harvested cells were resuspended in fresh complex culture medium containing 5 mM MV, 5 mM DT. Substrates final concentration was 20 mM. The reaction mixture was incubated at 40 °C for 15

19 Reduction of different substrates by pure NarGH (B). Assays were carried out in 20 50 mM Tris buffer pH 8 at 40 °C for 15 minutes. Substrates final concentration 21 was 30 mM. 100 % Activity = 0.28 μ moles MV oxidized min⁻¹ mg prot.⁻¹ MV assay 22 was used in this experiment.

minutes. 100 % Activity = 0.76 U.

Figure 2. Optimum pH (A) and effect of salts concentration on chlorate reductase
activity (B) Reactions mixtures were incubated at 40 °C for 10 minutes. In panel
A, 100 % Activity correspond to 0.13 μmoles KClO₃ reduced min⁻¹ mg prot.⁻¹. Otoluidine assay was used in this experiment.
Figure 3. Effect of temperature and salt concentration on chlorate reductase
activity. Assays carried out in 50 mM Tris buffer pH 8, at 40 °C, 10 minutes. Otoluidine assay was used in this experiment.

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Figure 4. Nitrate reductase activity in presence of different oxyanions. (triangle: 18 mM nitrate + bromate; rhombus: 18 mM nitrate + perchlorate; square: 18 mM nitrate + chlorate). MV was used in this assay as electron donor. Nitrite production by pNarGH was follow using Griess method.

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Figure 5. Protein sequence aligment *Hfx. mediterranei* chlorite dismutase-like protein with *Hfx. volcanii* PitA and *Halorubrum lacusprofundi* B9LRB6, *Haloterrigena turkmenica* D2RQG0 and *Natrialba magadii* D3STR5 using Clustal 2.0.12. Histidine residues lying in the region linking chlorite dismutase-like and monooxygenase-like domains are in bold and underlined.

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- **Figure 1**



В





- 1 Figure 3



Figure 4







2 Figure 5

3	н	mediterranei		48		
Ă	л т	wolgopii		10		
3	п 17	Voicanii		40		
S	Н	lacusprofundi	MVEAPQIEEGWFALHDFRSIDWDAWRDAPERERKRAIEEGKAFLKHRE	48		
g	Η	turkmenica	MERRQPPQTEEGWYVLHDFRSIDWDAWRDAPERRRSRAIEEGIEYLSAAE 5			
	Ν	magadii	MERRQPPQTDEGWYVLHDFRSIDWDAWRDAPEHRRSRALEEGIDYLTAAN	50		
ğ						
.9	Η	mediterranei	AVEDAAEGTSAIFSVLGHKADFMVVHFRPTLDDISRAERQFEQTALAEFT	98		
10	Η	volcanii	AVEDAEAGASAVFSVLGHKADFVVVHFRPTLDDISRAERQFERTALAAFT	98		
11	Η	lacusprofundi	LVADADEGDSGLFSVLGHKADLLFVHFRPTLDDLSSIERRFEDTALANFT	98		
12	Н	turkmenica	SVADAEEGDSATFAVLGHKADLLVLHLRPTLADLDALERRFEGTALAEFT	100		
13	N	magadii		100		
ĺΔ	14	magaarr		100		
15				140		
16	Н	medilerranei	EQPISION VEVSELVSDIFEGNKEDIDIGLERIEGKEQPDIPDDIM	148		
17	Н	volcanii	EQPTSYVSVTEVSGYVSDDYFEGNEEDIDAGLLRYIEGKLKPDIPEDTYM	148		
1/	Η	lacusprofundi	ERTTSYVSVTEVSGYVSDEFFE-DPESVDTGLKRYIEGKMTPEIPDDEYV	147		
18	Η	turkmenica	ERADSYLSVTEVSGYMSQDYFDEDAEVEDTGMARYIETRLKPEIPDSEFL	150		
19	Ν	magadii	ERADSYLSVTEVSGYMSQEYFEEDGEIEDTGTKRYIESRLKPTIPDSEFV	150		
20						
21	Η	mediterranei	SFYPMSKRRGEKHNWYDLPFDERRELMSVHGDTGRKYAGKIKQVIASSVG	198		
22	Η	volcanii	SFYPMSKRRGEEHNWYDLSFDERRDLMSTHGDTGROYAGKIKOVIASSVG	198		
$\overline{2}\overline{3}$	Н	lacusprofundi	CEVENSKERGEEVNWYDLSEEDRADLMADHGEVGKEVAGKIKOVIASSVG	197		
$\overline{2}\underline{4}$	и Ц	turkmenica		200		
55	11	mamadii	SFIFMDRRRGFEDIWIDDEFDERAEIIDSSIGDIGRDIAGRVIQIISGSIG	200		
25	11	Magadii	SF IPMDKRKGPEHNWIDLPFDERADHLSSHGELGRNIAGRVIQIISGSVG	200		
57				040		
56	Н	meaiterranei	FEEFEWGVTLFGDDPTDIKDIVYEMRFDEVSAKYGEFGEFYVGRRFPPSD	248		
20	Η	volcanii	FDDYEWGVTLFGDDPTDIKDIVYEMRFDEVSSKYGEFGQFYVGRRFPPSD	248		
29	Η	lacusprofundi	FDSHEWGVTLFGSDPTDIKDIVYEMRFDPASSRYGEFGEFYIGRRFPPED	247		
30	Η	turkmenica	LDDFEWGVTLFGDDPTDVKELLYEMRFDPSSSRFAEFGRFLSARRFPPED	250		
31	Ν	magadii	LDDFEWGVTLFADDPTDVKELLTEMRFDPSSSAFAEFGRFLSARRFPPAN	250		
32						
33	Η	mediterranei	LGAFLAGDAVPTSEFGDESHHHAHAHG-EGGHHHGEGGHAHGEDGH	293		
34	Н	volcanii	LGAFLAGDGVPTSEFGDESHHGAHAHG-EGG-HHGEGGDGH	287		
35	Н	lacusprofundi	LGAYFAGETVPTPAGDGDTGDTEDGHGHAHG-EGHDHAGSGGGSAHGD	294		
36	н	turkmenica		295		
ŽŽ	M	magadii		200		
38	11	magadii		290		
žõ				221		
10	Н	medilerranei	HHGESGHGHGEGGHHGGDSDDEADEIDIRGQLDDLNIY	331		
40	Н	Volcanii	HHHDDGDGDHPHGDDGDEAADEDIRGQLEDLNIY	321		
41	Н	lacusprofundi	HPHGEEETSGEGDHPHSGEEGGHGGEDGDDPSDAD1RGELADLN1Y	340		
42	Η	turkmenica	HHHGDSSSSGRGDHGGSGGPHGDDDEDLRSELEDMGVY	333		
43	Ν	magadii	HHHGDSG-HGHGHGHGSGDPHDDAGADEDDESVRSELEELGVY	332		
44						
45	Η	mediterranei	AGKPHGEDVYATVLYSEADADEVFEEVEGLRGNFDHYPTHVKTAVYEAND	381		
46	H	volcanii	AGKPHGEDVYATVLYSEADADELFEEVEGLRGNFDHYPTHVKTAVYEANE	371		
47	Η	lacusprofundi	AGKPSGEDVYATVLYSEADVDELFDEVEGLRGNFDHYGTHVKTAVYEGRV	390		
48	Н	turkmenica	AGOPHGEDVHAVVLYSAADAEELFEEVDGLRGNEDHYDTHVKTAVYEPOD	383		
4 <u>9</u>	N	magadii	AGOPHGEDVHAVVLYSAADAGELEEEVEGLETNEDHVDTHVKTAVVEDON	382		
50	±•			202		
5ĭ	ц	mediterranei		17F		
š 2	п 17	wolappii		110		
52	п		RGRVAVVSIWEIASAALIAAGFLSELPGIVERAGEGSGFGIMGMF	410		
55	Н	iacusproiundi	IDKAAVVSIWDTASAAETAAGFLSELPEVVARAGEESGFGTMGMF	435		
24	Η	turkmenica	GGDDSETAVVSLWETERAASTAAGFLADLPDIVRQAGDDEGDSWGTMGMF	433		
22	Ν	magadii	DDSDAETAVVSLWETDRAANTAAGFLADLPDIVRQAGDDEDDSWGTMGMF	432		
ŽĎ						
51	Η	mediterranei	$\verb YTVKPEHREDFVEKFGVVGGLLDDMDGHFDTDLMVNLEDENDMFIASQWR $	476		
58	Η	volcanii	YTVKSEHRGDFVEKFGTVGGLLEEMDGHFDTDLMVNVEDEDDMFIASQWR	466		
59	Η	lacusprofundi	YTVKPEHQEDFTDTFDDVGEILAEMDGHVETDLMMNVEDENDMFIASQWH	485		
60	Η	turkmenica	YSVKPEHRGDFLGTFEEAGELLAEMDGHRKTDLLINREDENDMFIASRWD	483		
61	N	magadii	YTVKPEHRGDFTGVFDDAASTLAEMDGHRKSDLUVNREDENDMFTASRWD	482		
Ğ2	±•			102		
63	ц	mediterranei				
64	11 17	wolaanii				
65	п					
66	H	racusprorunai	AREDAMAFFGODEFREIVUNGREVLADRPRHVFLA 520			
60	Н	turkmenica	SKEDAMQFFRSDAFSEAVEFGRDVLTDRPRHVFLA 518			
0/	Ν	magadii	SREDAMQFFRSDEFAETVEFGRDVLADRPRHVFLA 517			
68						

Table 1: Specific growth rate (μ) and cell doubling time (t_d) calculated from *Hfx. mediterranei* cultures anaerobically grown in complex media [20] in presence or
 absence of electron acceptor.

Electron acceptor Optical density		Batch culture time	μ (h⁻¹)	t _d
	(600nm) at	(hours)		(hours)
	stationary			
	phase of growth			
None	0.15	150	0.0058	79
Nitrate	2	150	0.11	6.5
Chlorate	0.15	150	0.0060	75
Nitrate growing	0.8	150	0.019	27
cells transferred to				
chlorate media				

Table 2: Turnover numbers (Kcat) and kinetic constants for NarGH. Kinetic parameters related to nitrate where obtained at pH 8 in presence of 3.6 M NaCl and MV as electron donor. Regarding to chlorate, kinetic parameters where obtained in presence of 0.5 M NaCl and MV as electron donor. Activity is expressed as μmoles MV oxidized min⁻¹ mg prot.⁻¹

Substrate	K _m	V_{max} (µmoles MV oxidized	K _{cat} (s ⁻¹)	K _{cat} / K _m	Reference
	(mM)	min ⁻¹ mg prot. ⁻¹)		(mM⁻¹s⁻¹)	
Nitrate	0.8	0.25	0.042	0.052	[20, 22]
Chlorate	2.4	0.28	46	19	This paper